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## Research Article

### Derivation of Point of Departure (PoD) Estimates in Genetic Toxicology Studies and Their Potential Applications in Risk Assessment

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Genetic toxicology data have traditionally been employed for qualitative, rather than quantitative evaluations of hazard. As a continuation of our earlier report that analyzed ethyl methanesulfonate (EMS) and methyl methanesulfonate (MMS) dose-response data (Gollapudi et al., 2013), here we present analyses of 1-ethyl-1-nitrosourea (ENU) and 1-methyl-1-nitrosourea (MNU) dose-response data and additional approaches for the determination of genetic toxicity point-of-departure (PoD) metrics. We previously described methods to determine the no-observed-genotoxic-effect-level (NOGEL), the

breakpoint-dose (BPD; previously named Td), and the benchmark dose (BMD<sub>10</sub>) for genetic toxicity endpoints. In this study we employed those methods, along with a new approach, to determine the non-linear slope-transition-dose (STD), and alternative methods to determine the BPD and BMD, for the analyses of nine ENU and 22 MNU datasets across a range of *in vitro* and *in vivo* endpoints. The NOGEL, BMDL<sub>10</sub> and BMDL<sub>1SD</sub> PoD metrics could be readily calculated for most gene mutation and chromosomal damage studies; however, BPDs and STDs could not always be derived due to

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data limitations and constraints of the underlying statistical methods. The BMDL<sub>10</sub> values were often lower than the other PoDs, and the distribution of BMDL<sub>10</sub> values produced the lowest median PoD. Our observations indicate that, among the methods investigated in this study, the BMD approach is the preferred PoD for quantitatively describing genetic toxicology

data. Once genetic toxicology PoDs are calculated via this approach, they can be used to derive reference doses and margin of exposure values that may be useful for evaluating human risk and regulatory decision making. *Environ. Mol. Mutagen.* 00:000–000, 2014. © 2014 Wiley Periodicals, Inc.

**Key words:** benchmark dose; ENU; MNU; alkylating agents; margin of exposure

## INTRODUCTION

Until quite recently genotoxicity test results were employed almost exclusively for dichotomous qualitative evaluations (i.e., results classified as either a positive or negative), with studies routinely evaluating responses at very high doses (i.e., near the maximum tolerated dose or MTD). Interest in the manifestation of genotoxicity at low doses, as well as quantitative analyses of the dose–response data, has been limited (Pottenger and Gollapudi, 2009). In contrast, quantitative dose–response analyses and derivation of point-of-departure (PoD) metrics are routinely employed to assess other toxic effects that are not mediated by genotoxic mechanisms (Piersma et al., 2011) as well as carcinogenic risk (EFSA, 2009). Such PoD values are routinely used for risk assessment in conjunction with uncertainty factors to derive health-based guidance values and regulatory limits to assess and manage risks. In the case of genotoxic carcinogens, if the mode of action is not established, then a conservative, linear approach (i.e., linear low-dose extrapolation from the PoD to origin) is generally taken. Thus, there is theoretically no level of exposure for such a chemical that does not pose a small, but finite probability of generating a carcinogenic response. This assumption may not always hold true, because there is increasing recognition that non-linear dose responses are observed with genotoxic endpoints for at least some substances, and there is increasingly strong mechanistic evidence to support the calculation and use of biologically meaningful PoDs to inform regulatory decision making for genotoxic agents. For example, several recent publications have demonstrated that biologically meaningful, sub-linear dose–response functions exist for both non-DNA-reactive genotoxins [mitotic spindle poisons (Johnson and Parry, 2008; Elhajouji et al., 2011)] and at least some DNA-reactive mutagens (Doak et al., 2007; Gocke and Wall, 2009; Johnson et al., 2009; Pottenger et al., 2009; Bryce et al., 2010; Gollapudi et al., 2013). This recognition has contributed to an increasing appreciation of the utility of the quantitative analysis of genetic toxicity dose–response relationships; and moreover, to employ quantitative methods and PoD determination for genotoxicity data to use in regulatory decision making.

The Quantitative Analysis Workgroup (QAW) of the Genetic Toxicology Technical Committee (GTTC) coordi-

nated by the Health and Environmental Sciences Institute (HESI) of the International Life Sciences Institute (ILSI) is involved in the development and critical examination of methodologies for the quantitative analysis of *in vitro* and *in vivo* genotoxicity dose–response data, and the development of strategies for the use of PoD metrics to support regulatory evaluations and decision making (Gollapudi et al., 2011). In this report, the GTTC QAW extends the analyses presented earlier (Gollapudi et al., 2013) that addressed the applicability of several statistical methods for the analyses of genetic toxicology dose–response data. Collectively, this and our previous report contribute to a rapidly growing body of knowledge regarding the use of quantitative dose–response analyses to derive PoD metrics, and moreover, to employ PoD values to assess the risk of adverse health effects in humans and/or to determine exposure levels that would be associated with negligible risk.

Gollapudi et al. (2013) used data from studies of methyl methanesulfonate (MMS) and ethyl methanesulfonate (EMS) to investigate the utility of several metrics to define a PoD for use in determining regulatory limits associated with negligible risk of genotoxic effect. These metrics included (i) no-observed-genotoxic-effect-levels (NOGELs), (ii) statistically defined break points, now referred to as the breakpoint dose (BPD, previously termed “threshold” dose, T<sub>d</sub>), and (iii) benchmark dose (BMD) levels (Gollapudi et al., 2013). This report addresses the need to extend this initial investigation to encompass additional agents, to critically examine additional modeling techniques, to improve the biological understanding of the mode(s)-of-action (MOAs) that determine(s) the shape(s) of genotoxicity exposure-response curves, and to define strategies to employ PoD metrics for regulatory evaluations and decision making. Such work is essential, and will precede general acceptance regarding the determination and use of meaningful PoD metrics in genetic toxicology. The derivation of PoDs in genetic toxicology studies and their routine use for risk assessment and regulatory decision making will require the following:

1. Selection of appropriate mathematical models and statistical methods for reliable PoD determinations;
2. Determination of study design features that facilitate quantitative dose–response analyses, and,

3. Development of standardized methods for the incorporation of PoD metrics into human health risk assessment (e.g., application of uncertainty and/or safety factors and/or margin of exposure analysis to define exposures that are associated with negligible risk).

The current work extends our earlier efforts to address points 1, and 2 above, and to initiate discussions on point 3. This was carried out using the data on two potent alkylating mutagens with large data bases, 1-methyl-1-nitrosourea (MNU) and 1-ethyl-1-nitrosourea (ENU) and, building on our earlier analyses of the related, but less potent alkylating agents EMS and MMS, enabling further development of our “toolbox” for the derivation and use of PoD metrics in genetic toxicology.

## MATERIALS AND METHODS

### Data Selection

The database employed, referred to as the G4 database, was developed by the GTTC QAW, and details of the quality criteria used for data screening are in Gollapudi et al. (2013). In addition to the EMS and MMS datasets, the database contains a total of 45 datasets for ENU and MNU, including endpoints for gene mutation and chromosomal damage (measured as micronucleus formation) *in vitro* and *in vivo*. Since the goal of this work was to analyze datasets from which four PoD values could be derived (i.e., NOGEL, BMD, BPD, and STD), only datasets with data supporting PoD derivation were evaluated. Based on the recommendations of Lutz and Lutz (2009), we further restricted the analyses to datasets with  $\geq 5$  doses (including the negative control) to ensure that we could use the bilinear modeling approach (i.e., BPD modeling).

### Benchmark Dose Analysis

The benchmark dose (BMD) is defined in the U.S. Environmental Protection Agency's (EPA's) risk assessment glossary as 'A dose or concentration that produces a predetermined change in response rate of an adverse effect (called the benchmark response or BMR) compared to background' (<http://www.epa.gov/riskassessment/glossary.htm>). For BMD determination, the latest versions of the PROAST BMD software and EPA's Benchmark Dose Software (BMDS) were employed (EPA, 2013).

### RIVM-PROAST Benchmark Dose Analysis

BMD analysis was conducted using PROAST, the dose-response modeling software developed at the National Institute for Public Health and the Environment (RIVM) in The Netherlands [<http://www.proast.nl>; version 36.9 (Slob, 2002)]. The nested set of models used included the exponential and Hill models that are recommended by the European Food Safety Authority (EFSA, 2009). Note that the bilinear models used to define BPD metrics are not included, and thus are not assessed in these BMD packages (Slob and Setzer, 2013). The BMR examined was 10%, which corresponds to an increase equal to 10% of the background (negative control) level, as estimated by the fitted model used for continuous endpoints such as genotoxicity. This is an arbitrary choice, but is one that has been frequently employed in the literature and that provides a conservative PoD that is close enough to the observable range of dose-response data to allow reliable estimates to be derived from fitted curves without uncertain extrapolation. A BMR of 10% is also the approach currently being used in standard toxicology dose-response

analyses. This allows a better comparison of the genotoxicity data to the other toxicity data when performing a risk characterization for human health. This BMR that corresponds to an increase above the background, differs from the 10% increase in incidence (e.g., the additional risk or extra risk) used for quantal data such as carcinogenicity data. The lower limit of the one-sided 90% confidence interval on the BMD is termed the BMDL, and the BMDL<sub>10</sub> refers to the estimate of the lower 90% confidence limit of a dose that produces a 10% increase over the fitted background level for continuous endpoints. The BMDU<sub>10</sub> is the upper limit of this 90% confidence interval. Model selection was performed based on a log-likelihood ratio test that assesses whether including additional parameters to the model results in a statistically significant improvement in model fit (Hernández et al., 2011a). The model with additional parameters is accepted only if the difference in log-likelihoods exceeds the critical value at  $P = 0.05$ . In addition, the log-likelihood is used to compare the “full” model (geometric means of the observations at each dose) to the selected model, to provide an indication of the goodness-of-fit. The distribution of the residual errors in PROAST is similar to that derived using the EPA's BMDS (discussed below). For continuous data, the residual errors are assumed to be log-normally or normally distributed. There is an option in both software packages to choose one or the other distribution. In PROAST, the default assumption is that the standard deviation is proportional to the mean, and thus a log-normal distribution is applied to the continuous data, whereas in BMDS the default setting is the normal distribution.

### U.S. EPA Benchmark Dose Software

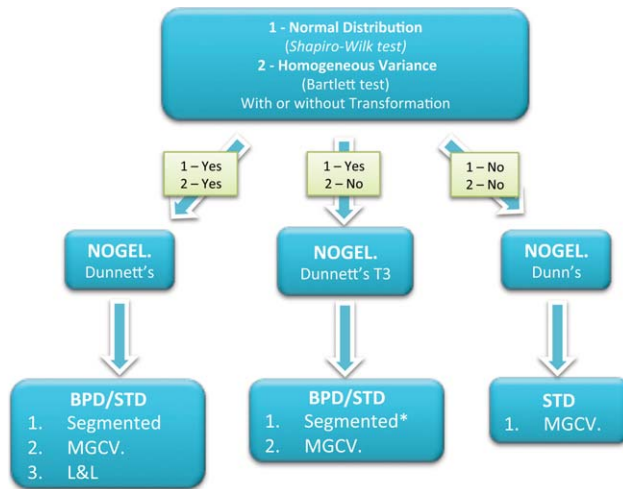
BMD analysis also was conducted using the latest version of the EPA's Benchmark Dose Software [i.e., BMDS v2.4 (EPA, 2013)]. The standard suite of continuous models (Hill, exponential, polynomial, linear, and power) was used along with constant and non-constant variance model assumptions. As was the case for RIVM-PROAST, the bilinear models used to define BPD metrics are not included, and thus are not assessed in this BMD package. The BMR chosen for the BMDS software was an increase equivalent to one standard deviation above the spontaneous (control) value; this was used in the calculation of a one-sided 95% lower confidence limit for this BMD value and designated BMDL<sub>1SD</sub>. This BMR is recognized to be equivalent to  $\sim 10\%$  excess risk for individuals below and above the 2nd and 98th percentiles, respectively (Crump, 1995). Model selection was primarily based upon the  $P$ -value for goodness-of-fit to the data and the Akaike's Information Criterion (AIC). For this study, the best fitting model was selected among the suite of continuous models (see above). Log-transformation was used as a default for analysis of continuous data with PROAST (Slob and Setzer, 2013). For BMDS, continuous data were transformed in the same manner used for NOGEL, BPD, and smoothing regression spline modeling (discussed in next section).

### NOGEL, BPD, and Smoothing Regression Spline Analyses

NOGEL, BPD, and Smoothing Regression Spline analyses were used to extend the previous effort (Gollapudi et al., 2013) to examine the utility of open source methodologies and additional approaches.

### Initial Statistical Evaluation

All data sets were imported into R (R Development Core Team, 2011) and the following analyses conducted: Shapiro-Wilk Normality Test, Bartlett Test of Homogeneity of Variances, Jonckheere-Terpstra Test of Monotonic Trend (asymptotic version), and a Bonferroni test for outlier identification. Data were transformed in order to achieve normally distributed data and homogeneity of the within dose variances. In most cases where the original data were not normally distributed and/or the variances were heterogeneous, one of the data transformation processes (square root or logarithmic) performed on the response data



**Fig. 1.** Flow chart showing the workflow for determination of NOGELs, BPDs and STDs. In cases where the data do not meet parametric screening requirements even following variable transformation, models such as the segmented (see <http://cran.r-project.org/web/packages/segmented/segmented.pdf>)\* or mgcv (Mixed GAM [generalized additive model] Computation Vehicle, see <http://cran.r-project.org/web/packages/mgcv/mgcv.pdf>) can be utilized since they are less influenced by distributional problems. \*segmented can be used with T3, but requires special implementation not conducted in this study. All statistical approaches presented in this flow chart are now available for download and use in R (versions 3.0.2 and above), the ILSI-HESI GTTC QAW developed and successfully submitted the package called 'drsmooth' to CRAN in 2013 (Hixon and Bichteler, 2013). NOGEL, no observed genotoxic effect level; BPD, breakpoint dose; STD, slope transition dose; L&L, Lutz, and Lutz, 2009. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]

resulted in datasets with satisfactory distributions and variances, and the transformation essentially eliminated any significant outliers identified by the Bonferroni test. If transformation of data was necessary for a dataset, then all remaining analyses on that dataset were conducted on similarly transformed data. Based on these preliminary statistical tests, and using a criterion of  $P < 0.05$  to identify deviations from normality and heterogeneous variances, different pathways of statistical analyses and modeling approaches were used to derive NOGEL, BPD, and slope transition dose (STD) values (Fig. 1).

### No Observed Genotoxic Effect Level (NOGEL)

The NOGEL was defined as the highest tested dose at which there was no statistically significant increase in genotoxic effect compared to the control. Means from datasets that met the criteria for being normally distributed with homogeneous variances across dose groups were compared using a *post-hoc* Dunnett's Test ( $\alpha = 0.05$ ). Means for datasets that were normally distributed but with heterogeneous variances were compared using a *post-hoc* Dunnett's T3 test (Field, 2009). Means from datasets that were non-normally distributed and with heterogeneous variances were compared using the non-parametric *post-hoc* Dunn's Test (Laws et al., 2000).

### Breakpoint Dose (BPD) Modeling

Two similar bilinear methodologies were used to identify a discontinuity or breakpoint (below which the slope was zero) in the dose-response function: Lutz and Lutz (2009), and a package in the R system called 'segmented' (Muggeo, 2008; Wood, 2011). The Lutz and Lutz

model, referred to here as the L&L model, is the bilinear approach discussed and applied by Gollapudi et al., (2013). The R segmented package (Muggeo, 2008) is similar to the L&L model—i.e., it determines the best-fitting two segment linear function where the first segment from zero dose to the breakpoint is horizontal (i.e., has zero slope) and the second segment has a positive slope. However, the segmented approach has several advantages: it is based on an open source, peer reviewed package available in R, and, unlike the L&L model, it does not require removal of top doses due to supra-linearity (saturation or high-dose toxicity). With 'segmented', the breakpoint where the slope changes is called the BPD, with its standard error serving as the basis for the calculation of confidence bounds (i.e., 90% two-sided confidence bounds corresponding to the output of PROAST, BMDS, and the L&L models). In both the L&L and segmented models, a  $BPDL \leq 0$  indicates that a model with a single, linear segment with non-zero slope (i.e., a linear dose response) provides a better fit for the data. The L&L model relies on the assumption that the data are normally distributed and have homogeneous variances, and therefore was only used on data where the data transformation was successful. Although 'segmented' can be used in conjunction with weights to account for variance heterogeneity across dose groups, this approach was not used for analyses reported here. Therefore, 'segmented' was also only used on normally distributed datasets with homogeneous variances.

### Smoothing Regression Spline

We also used penalized smoothing splines to analyze the dose-response relationships. Penalized smoothing splines are a family of flexible techniques for estimating a continuous functional relationship without the need to assume linearity or any specific non-linear functional form. Wood (2006) has extensively developed the underlying theory. The 'mgcv' (Mixed GAM [generalized additive model] Computation Vehicle) package in R (Wood, 2006, 2011) was used to estimate the dose-response function using a default thin plate smoothing regression spline with degrees of freedom determined by generalized cross validation. Unlike the bilinear model used with the 'segmented' algorithm, 'mgcv' is commonly regarded as semi-parametric because it can be applied to non-normally distributed data with heterogeneous variances, and still provide an optimal solution from a cross-validation perspective. Thus, smoothing spline regression was used for PoD determination irrespective of whether the data required a parametric or non-parametric test (see Fig. 1).

The derived continuous non-linear dose-response function was subjected to finite differencing to calculate its first derivative, or slope, along the length of the function. The standard error for the slope also was calculated and used to form confidence limits. These served as the basis for the determination of the slope transition dose, or STD. The STD given by the smoothing regression spline is the lowest dose at which the dose-response function has a slope that is significantly above zero with 95% confidence using a one-sided confidence limit. As above, this particular limit was chosen to match the output of PROAST, BMDS, and the L&L model. At all lower doses, the slope of the dose-response function is not statistically distinguishable from zero, therefore represents a flat line. When the lower bound CI on the STD (i.e.,  $STD_L$ ) is  $\leq 0$ , the hypothesis that the slope is increasing significantly at dose = 0 (i.e., slope is above zero) cannot be rejected. This is conceptually different from the L&L and segmented bilinear models, where a  $BPDL \leq 0$  means that linearity cannot be rejected (regardless of the slope of that linearity). The smoothing regression spline approach arrives at a PoD by directly assessing the slope of the dose-response relationship throughout the dose range to determine the dose where it initially becomes positive.

In addition to deriving potential PoDs, the smoothing regression spline model also was used to test the overall linearity of the dose response of a given dataset. To this end, we tested whether the smoothing regression spline model fit the overall dose response significantly

**TABLE I. Summary of Datasets Analyzed, Including Information on Endpoint, Type of Experimental System, Tissue or Cell Type, Treatment Regime, Dose Units, and Source**

Chemical	Endpoint	Gene target	Type	Species	Cell type/tissue	Units	# of doses	# of replicates	Treatment regime	Expression time <sup>a</sup>	Study
ENU	Gene mutation	<i>Dlb1</i>	<i>vv</i>	Mouse	SI	mg/kg	5	3-4	1 d/i.p.	49 d	van Delft et al. (1998)
	Gene mutation	<i>LacZ</i>	<i>vv</i>	Mouse	SI	mg/kg	5	4	1 d/i.p.	49 d	van Delft et al. (1998)
	Gene mutation	<i>LacZ</i>	<i>vv</i>	Mouse	Spleen	mg/kg	5	4	1 d/i.p.	49 d	van Delft et al. (1998)
	Gene mutation	<i>Pig-a</i>	<i>vv</i>	Mouse	RET	mg/kg	7	5	1 d/i.p.	14 d	Bhali et al. (2011)
	Gene mutation	<i>Pig-a</i>	<i>vv</i>	Mouse	RBC	mg/kg	7	5	1 d/i.p.	28 d	Bhali et al. (2011)
	Gene mutation	<i>HPRT</i>	<i>vt</i>	Human	AHH-1	µg/mL	12	3	24 hr	13 d	Doak et al. (2007)
	Micronucleus			Mouse	RET	mg/kg	7	5	1 d/i.p.	24 hr	Bhali et al. (2011)
	Micronucleus		<i>vt</i>	Human	TK6	µg/mL	18	4	24-30 hr	0 hr	Bryce et al. (2010)
	Micronucleus		<i>vt</i>	Human	AHH-1	µg/mL	15	2-3	18 hr	0 hr	Doak et al. (2007)
	Gene mutation	<i>Hprt</i>	<i>vv</i>	Mouse	Spleen	mg/kg	5	9-10	1 d/i.p.	21 d	Monroe et al. (1998)
MNU	Gene mutation	<i>LacI</i>	<i>vv</i>	Mouse	Spleen	mg/kg	5	6-7	1 d/i.p.	21 d	Monroe et al. (1998)
	Gene mutation	<i>Pig-a</i>	<i>vv</i>	Rat	RET	mg/kg	7	3	28 d/gavage	15 d	Lynch et al. (2011) <sup>b</sup>
	Gene mutation	<i>Pig-a</i>	<i>vv</i>	Rat	RBC	mg/kg	7	3	28 d/gavage	29 d	Lynch et al. (2011) <sup>b</sup>
	Gene mutation	<i>Pig-a</i>	<i>vv</i>	Rat	RBC	mg/kg	5	6	28 d/gavage	4, 15, 29 d	BMS (unpublished) <sup>c</sup>
	Gene mutation	<i>Pig-a</i>	<i>vv</i>	Rat	RET	mg/kg	5	6	28 d/gavage	4, 15, 29 d	BMS (unpublished) <sup>c</sup>
	Gene mutation	<i>HPRT</i>	<i>vt</i>	Human	AHH-1	µg/mL	11	3-4	24 hr	13 d	Doak et al. (2007)
	Gene mutation	<i>Tk</i>	<i>vt</i>	Mice	L5178Y	µM	10	5	4 hr	2 d	Pottenger et al. (2009)
	Gene mutation	<i>HPRT</i>	<i>vt</i>	Human	AHH-1	µg/mL	10	3	24 hr	13 d	Thomas et al. (2013)
	Micronucleus		<i>vv</i>	Rat	PCE	mg/kg	8	6	4 d/gavage	24 hr	LeBaron (2009)
	Micronucleus		<i>vv</i>	Rat	NCE	mg/kg	8	6	4 d/gavage	24 hr	LeBaron (2009)
Micronucleus	Micronucleus		<i>vv</i>	Rat	RET	mg/kg	7	3	28 d/gavage	4, 29 d	Lynch et al. (2011) <sup>b</sup>
	Micronucleus		<i>vv</i>	Rat	RBC	mg/kg	7	3	28 d/gavage	4, 29 d	Lynch et al. (2011) <sup>b</sup>
	Micronucleus		<i>vv</i>	Rat	RET	mg/kg	5	6	28 d/gavage	4, 29 d	BMS (unpublished)
	Micronucleus		<i>vt</i>	Human	TK6	µg/mL	18	4	24-30 hr	0 hr	Bryce et al. (2010)
	Micronucleus		<i>vt</i>	Human	AHH-1	µg/mL	9	4	18 hr	0 hr	Doak et al. (2007)

*vv*, *in vivo*; *vt*, *in vitro*; RET, reticulocytes; RBC, red blood cells; NCE, non-chromatic erythrocytes; PCE, polychromatic erythrocytes; IP, intraperitoneal injection; SI, small intestine; d, day; hr, hour.

<sup>a</sup>Assume post treatment unless multiple time points are listed, indicating time after initial dosing.

<sup>b</sup>Study 2 from Lynch et al. (2011).

<sup>c</sup>BMS unpublished, Bristol-Myers Squibb unpublished data provided to HESI for incorporation into the G4 database.

**TABLE II. Point of Departure (PoD) Metrics for ENU (1-Ethyl-1-Nitrosurea)**

Study	Type	Endpoint	Response Transformation	Trend Test	Slope< NOGEL	Linear< NOGEL Test	NOGEL	L&L BPD	mgcv STD	BMDU <sub>10</sub> <sup>a</sup>		
										BMDS	BMDL <sub>10</sub>	PROAST
VanDelft (1998) <i>Dlb1</i> _SI Mouse	vv	GM	LogR	7.60E-07	NA	NA	Dunnett's	NA	no BPD	1.55	0.09	18.05 mg/kg
VanDelft (1998) <i>LacZ</i> _SI Mouse	vv	GM	Raw	1.20E-06	0	ID	Dunnett's	10	no BPD	5.46	1.94	5.20 mg/kg
VanDelft (1998) <i>LacZ</i> _Spleen	vv	GM	Raw	3.80E-05	0	ID	Dunnett's	25	no BPD	6.00	11.67	1.38 mg/kg
Mouse												
Bhalai (2011) <i>Pig-a</i> _PCE Mouse	vv	GM	SqrR	8.30E-12	+	ID	Dunnett's	10	no BPD	no STD	1.46	0.33
Bhalai (2011) <i>Pig-a</i> _RBC Mouse	vv	GM	LogR	4.50E-12	+	ID	Dunnett's	45	NA	no STD	0.95 <sup>a</sup>	0.12
Bhalai (2011) PCE Mouse	vv	GM	LogR	0.0006	+	ID	Dunnett's	10	no BPD	no STD	4.03 <sup>a</sup>	1.36
Doak (2007) AHH1_Human HPRT	vt	GM	Raw	2.20E-15	0	ID	Dunnett's	0.2	0.57	no STD	0.60	0.18
Bryce (2010) TK6_Human Expt 1	vt	MN	LogR <sup>b</sup>	9.50E-08	0	no	Dunn's	2.34	NA	NA	1.38 <sup>a</sup>	0.84
Bryce (2010) TK6_Human Expt 2	vt	MN	LogR <sup>b</sup>	0.03	0	no	Dunnett's	9.37	5.24	3.86	5.95	8.07 <sup>a</sup>
Doak (2007) AHH1_Human	vt	MN	Raw	9.30E-08	+	yes	Dunnett's	0.4	no BPD	no STD	0.08	5.02

vv, *in vivo*; vt, *in vitro*; NA, not applicable; ID, insufficient doses; GM, gene mutation; MN, micronucleus; SI, small intestine; +, positive gradient; NOGEL, no observed genotoxic effect level; BPD, breakpoint dose; BPDL, breakpoint dose lower confidence interval; STD, slope transition dose; STDL, slope transition dose lower confidence interval; BMDL<sub>10</sub>, benchmark dose 10 lower deviation lower confidence interval; BMDL<sub>10</sub>, benchmark dose 10 upper confidence interval; L&L, Lutz and Lutz 2009.

<sup>a</sup>Poor fit for benchmark dose model,  $P < 0.05$ .

<sup>b</sup>Doses log transformed as well.

Response Transformation, same number added to 'R' to ensure all responses were above the value of 1 before transformation with Log or Sqrt.

'Slope<NOGEL' tests whether slope up to and including the NOGEL is fit better by linear or nonlinear model (i.e., smoothing regression spline).

better than a linear model. Therefore, if the derived model indicated nonlinearity, the shape of the dose response at the lower doses also was assessed, specifically the experimental doses below the lowest-observed-genotoxic-effect-level (LOGEL) (i.e., from zero dose, up to and including the NOGEL). This approach builds on the previous study, in which similar prerequisites were defined for the bilinear model (Gollapudi et al., 2013). Two tests were conducted. First, a simple test was used to determine whether the slope from zero dose to the NOGEL was significantly different from zero. The second test, which requires more doses than the first, determined whether the smoothing regression spline model fits the data from zero dose to the NOGEL significantly better than a linear model. This approach has been developed into a package called 'drsmooth', which is available for free download and use with R versions 3.0.2 and above (Hixon and Bichteler, 2013).

## RESULTS

Table I summarizes the individual study characteristics for the various genotoxicity datasets analyzed. A wide range of endpoints were analyzed, including *in vivo* and *in vitro* measures of micronuclei (MN), in various cell types and species. Similarly, gene mutations were assessed in multiple target genes including transgenes (e.g., *LacZ*) and endogenous genes (e.g., *Hprt*).

The PoD metrics for nine ENU genotoxicity datasets are summarized in Table II. NOGEL values were obtained for all datasets with the exception of the *Dlb1* mutation analyses in the small intestine, where the lowest study dose was significantly different from control. BMDL<sub>10</sub> values were determined using PROAST for reasons stated in Gollapudi et al. (2013), whereas BMDL<sub>10</sub> values were determined using BMDS. The latter is the default metric used by the EPA for continuous data (U.S. EPA, 2012). In all cases, an exponential model provided the best fit in PROAST. In BMDS, the best fitting model was selected among the typical suite of continuous models. Overall, although not all PoD methods provided a good fit of the data to a statistical model, the two BMD modeling approaches provided estimates of the BMDL values that were lower than the corresponding NOGELS.

PoD values were derived for four of the ENU datasets using at least one of the three models; L&L, segmented, and smoothing regression spline. In all four cases, the slope below the LOGEL was not significantly different from zero, suggesting a bilinear dose response. Two of the datasets for which BPD values were derived were not predicted to have an STD by mgcv (van Delft et al., 1998; Doak et al., 2007). Examination of the mgcv plots of these datasets reveals an apparent lack of bi-linearity, consistent with the model results failing to identify an STD (Table II). In contrast, the two datasets from Bryce et al. (2010) were predicted to have STD values generated by smoothing regression spline analyses (i.e. exhibited non-linear dose responses with estimated STDL values).

The PoD metrics for 22 MNU genotoxicity datasets are summarized in Table III. As with ENU, the two BMD modeling approaches yielded lower PoD values than the

TABLE III. Point of Departure (PoD) Metrics for MNU (1-Methyl-1-Nitrosourea)

Study	Type	Endpoint	Response transformation	Trend test	Slope < NOGEL	Linear < NOGEL Test	NOGEL	L&L	mgev segmented BPDL	BMDS BMDS <sub>ISD</sub>	PROAST BMDS <sub>ISD</sub>	PROAST BMDS <sub>10</sub>	PROAST BMDS <sub>10</sub> /BMDS <sub>10</sub> Units
Monroe (1998) HPRT_Spleen Mouse	vt	GM	LogR	2.80E+13	0	ID	Dunnett's	5	0.78	1.40	0.98	3.90	2.06
Monroe (1998) LacI_Spleen Mouse	vt	GM	LogR	0.005	0	yes	Dunnett's	15	8.22	no STD	10.62	11.53	5.26
Lynch (2011) <i>Pig-a</i> _RET Rat	vt	GM	Raw	0.048	NA	NA	Dunnett's	2.5	no BPD	no STD	no BPD	0.77	0.10
Lynch (2011) <i>Pig-a</i> _RBC Rat	vt	GM	SqrR	0.02	0	yes	Dunnett's	1.25	no BPD	no STD	0.19	0.77	0.11
BMS <i>Pig-a</i> _RBC Rat Day 4	vt	GM	LogR	0.26	NA	NA	Dunnett's	5	NA	no STD	NA	3.75	No DR
BMS <i>Pig-a</i> _RBC Rat Day 15	vt	GM	LogR	3.00E-06	+	yes	Dunnett's	2.5	NA	no STD	NA	1.57	0.20
BMS <i>Pig-a</i> _RBC Rat Day 29	vt	GM	LogR	8.50E-10	NA	NA	Dunnett's	None	no BPD	no STD	no BPD	0.43	0.015
BMS <i>Pig-a</i> _RET Rat Day 4	vt	GM	LogR	0.53	NA	NA	Dunnett's	5	NA	no STD	NA	4.33	No DR
BMS <i>Pig-a</i> _RET Rat Day 15	vt	GM	LogR	1.70E-07	0	ID	Dunnett's	0.9	no BPD	no STD	no BPD	0.74 <sup>a</sup>	0.10
BMS <i>Pig-a</i> _RET Rat Day 29	vt	GM	LogR	3.50E-08	NA	NA	Dunnett's	None	no BPD	no STD	no BPD	0.69	0.0007
LeBaron (2009) PCE_Rat	vt	MIN	LogR <sup>b</sup>	9.20E-09	+	yes	Dunnett's	1	NA	0.60	NA	0.08	0.02
LeBaron (2009) NCE_Rat	vt	MIN	LogR <sup>b</sup>	0.75	NA	NA	Dunnett's	50	NA	no STD	NA	42.2 <sup>a</sup>	No DR
Lynch (2011) PCE_Event 1 Rat	vt	MIN	Raw	4.50E-06	+	yes	Dunnett's	0.9	0.20	no STD	0.27	0.16	0.10
Lynch (2011) PCE_Event 2 Rat	vt	MIN	LogR	6.10E-07	+	yes	Dunnett's	0.6	0.15	no STD	no BPD	0.16	0.13
Lynch (2011) NCE_Event 1 Rat	vt	MIN	Raw	0.36	NA	NA	Dunnett's	2.5	no BPD	1.17	0.30	1.30	0.73
Lynch (2011) NCE_Event 2 Rat	vt	MIN	Raw	0.002	+	yes	Dunnett's	1.25	no BPD	no STD	no BPD	0.43	0.13
BMS Day 4 Rat	vt	MIN	LogR	8.80E-09	+	ID	Dunnett's	1.25	NA	0.56	NA	0.42	0.30
BMS Day 29 Rat	vt	MIN	LogR	3.60E-09	NA	NA	Dunnett's	None	no BPD	no STD	no BPD	0.40	0.18
Doak (2007) AHH1_Human	vt	GM	SqrR	1.00E-14	+	ID	Dunnett's	0.005	no BPD	no STD	no BPD	0.0004	0.0006
HPRT													1.51
Pottenger (2009) L5178Y_	vt	GM	LogR <sup>b</sup>	2.00E-08	0	yes	Dunnett's	0.69	1.01	0.49	1.03	0.83	0.61
Mouse Tk													2.03
Thomas (2013) AHH1_Human	vt	GM	SqrR	0.004	0	no	Dunnett's	0.0075	0.002	no STD	no BPD	0.008 <sup>a</sup>	0.006
HPRT													1.33
Bryce (2010) TK6_Human_Expt 1	vt	MIN	LogR <sup>b</sup>	0	+	no	T3	1.25	NA	0.07	0.08	0.47	0.26
Bryce (2010) TK6_Human_Expt 2	vt	MIN	LogR <sup>b</sup>	4.50E-14	+	no	Dunn's	0.23	NA	no STD	NA	0.20	0.066
Doak (2007) AHH1_Human	vt	MIN	Raw	2.50E-09	+	yes	Dunnett's	0.025	no BPD	no STD	no BPD	0.008	0.003

vt, *in vivo*; vt, = *in vitro*; NA, not applicable; ID, insufficient doses; GM, gene mutation; MN, micronucleus; No DR, no dose response; BMS, Bristol-Myers Squibb unpublished data; SI, small intestine; +, positive gradient; NOGEL, no observed genotoxic effect level; BPD, breakpoint dose; BPDL, breakpoint dose lower confidence interval; STD, slope transition dose; STDL, slope transition upper confidence interval; L&L, Lutz and Lutz, 2009.

Underlined PoD values were obtained after dropping high dose(s).

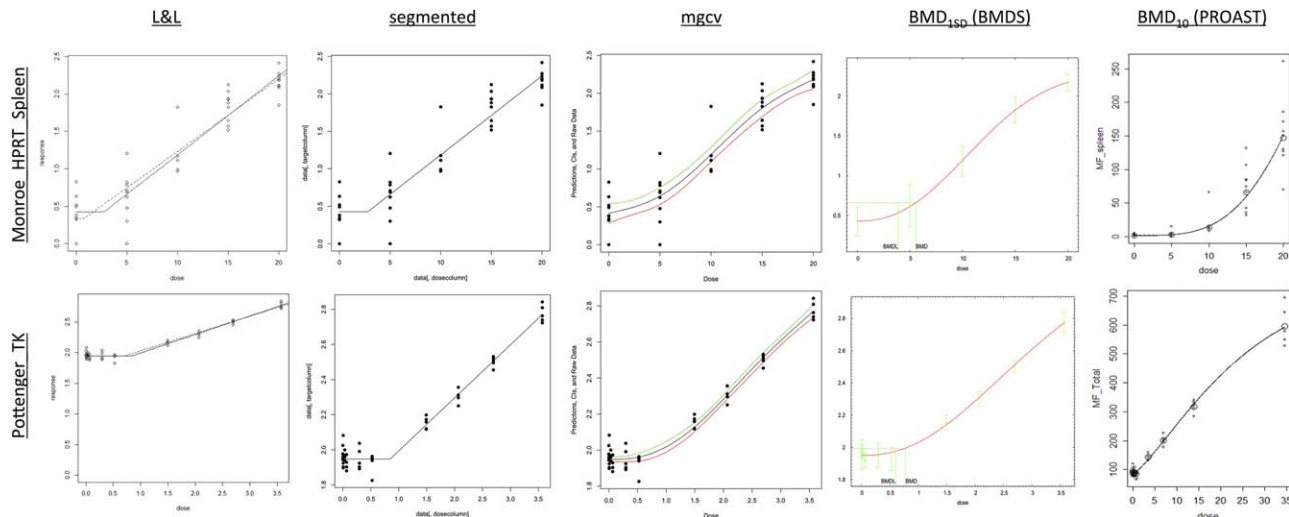
<sup>a</sup>Poor fit for benchmark dose model,  $P < 0.05$ .

<sup>b</sup>Doses log transformed as well.

Response Transformation, same number added to 'R' to ensure all responses were above the value of 1 before transformation with Log or Sqr.

'Slope < NOGEL' tests whether slope up to and including the NOGEL differs significantly from zero.

'Linear < NOGEL' tests whether slope up to and including the NOGEL is fit better by linear or nonlinear model (i.e., smoothing regression spline).



**Fig. 2.** Example of BPD, STD and BMD modeling results for two MNU gene mutation datasets (see results in Table III). The various software packages employed have only limited ability to adjust X- and Y-axes, and all plots shown are the default outputs of these programs. By default, PROAST provides plots with untransformed data. For both the Monroe and Pottenger datasets, log transformed responses provided the closest distribution to normal with homogeneous variance (see Table III).

L&L, Lutz, and Lutz hockey stick/bilinear approach for defining a breakpoint dose (BPD); Segmented, bilinear model for defining a BPD; mgcv, smoothing regression spline for defining a slope transition dose (STD) using 'drsmooth' in R version 3.0.2; BMD<sub>1SD</sub> (BMDS), Benchmark dose 1 standard deviation using BMDS; BMD<sub>10</sub> (PROAST), Benchmark dose 10 using PROAST. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]

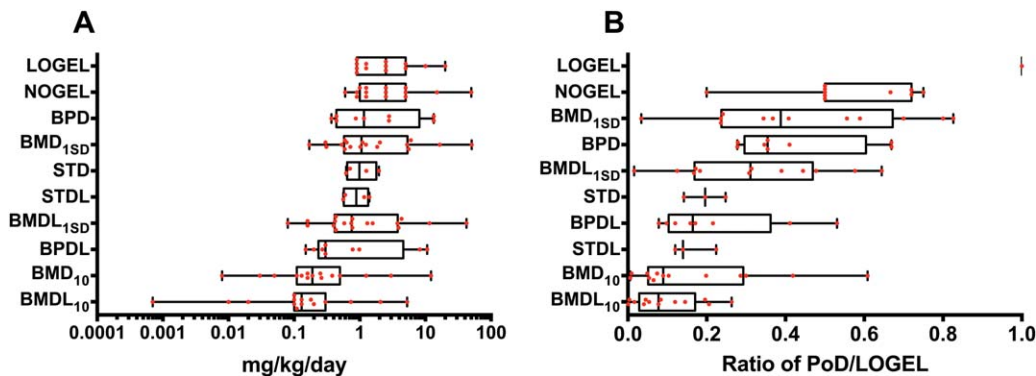
NOGEL. There were two MNU datasets for which both the bilinear and smoothing regression spline methods provided BPDLs and STDLs, respectively (Monroe et al., 1998; Pottenger et al., 2009). The PoDs associated with these data, calculated by the different aforementioned methods, were in remarkably close agreement for a given dataset, and the slopes up to and including the NOGEL were not significantly different from zero, demonstrating a good fit for the bilinear models (Fig. 2). There were eight datasets that required analysis with non-parametric methods, and the application of the smoothing regression spline methodology yielded STD values for five of these datasets, with the slopes from the negative control up to and including the NOGEL not being consistent with a zero slope. Therefore, these STD values should be interpreted with caution (see below).

Eighteen *in vivo* MNU datasets are summarized in Table III. Although the studies examined different endpoints in different species under different exposure scenarios, a quantitative comparison of the PoDs can still be conducted (Fig. 3). The medians (2.5 mg/kg/day) and distributions of NOGEL and LOGEL values were very similar (Fig. 3A). The BPD, STDL, and BMDL<sub>1SD</sub> were also in close agreement, ranging from 0.8 to 1.2 mg/kg/day. The median BPDL, BMD<sub>10</sub>, and BMDL<sub>10</sub> were 0.3, 0.2 and 0.1 mg/kg/day, respectively. To get a sense of how these PoD metrics compare to one another within each *in vivo* MNU dataset, each PoD was 'normalized' to the LOGEL of that dataset (if available). As expected, the ratio of the NOGEL to the LOGEL was less than unity (i.e., approximately 0.5; Fig. 3B). The ratios of the

BMDL<sub>1SD</sub> and BPD to LOGELs were similar, as were the ratios of STDL and BPDL to LOGELs. The ratio of the BMD<sub>1SD</sub> to the LOGEL was generally higher than the BPD and STD ratios, consistent with benchmark dose values being associated with a pre-defined increase in toxicological response. A similar pattern was observed for the BMDL<sub>1SD</sub> relative to the BPDL and STDL. The median ratios of the BMDL<sub>10</sub> values to the LOGELs were the lowest of all PoDs, and the BMDL<sub>10</sub> values were often lower than the BPDL and STDL ratios (Figs. 3A–B).

## DISCUSSION

The current work, which follows that of Gollapudi et al. (2013), is focused on advancing the development and application of statistical approaches to define PoDs for genotoxicity dose–response data. This is a necessary step in the path forward for the use of genotoxicity PoD metrics to inform regulatory decision making and/or risk assessment. The earlier work focused on the following PoD metrics: Dunnett's approach for calculating NOGELs; a multi-step approach to define BPD values (Gocke and Wall, 2009); and the BMD approach (i.e., using PROAST and BMDS software). Here we present the next phase in these analyses, which involved a series of approaches (Table IV) to determine PoDs for the potent genotoxins ENU and MNU. The data from the examined publications, in most cases, suggest that PoD values are generally lower and more difficult to define for ENU and MNU in comparison with EMS and MMS. This



**Fig. 3.** Comparison of PoD values for *in vivo* MNU genotoxicity datasets. (A) Box and whisker plots for PoDs listed in Table III (highest and lowest median values are at the top and bottom, respectively). (B) For each dataset, the PoD values (if available) were divided by the LOGEL for that dataset to provide the relative position of each PoD metric relative to the LOGEL (highest and lowest median ratios are at the top and bottom, respectively). The whiskers extend to min and max values, and the red dots represent individual values. There are fewer data points for certain PoD metrics, and the BPD values for segmented and L&L models

is likely because ENU and MNU are more potent mutagens, probably related to the higher proportion of  $O^6$ -alkylG and other pro-mutagenic adducts these  $S_N1$  alkylators form (Jenkins et al., 2005).

#### Evaluation and Comparison of Different Approaches for Determination of Point of Departure Values

Table IV presents a comparative assessment of the advantages and disadvantages associated with each PoD metric. The NOGEL value is the highest experimental exposure level at which there is no statistically significant increase above the concurrent experimental control value (background level). NOGELs are, by definition, dependent on study design features such as dose selection and the statistical power to detect an increase at each dose. Furthermore, this approach does not permit calculation of PoD confidence intervals. When comparing NOGELs to the other PoD metrics, one can see that NOGELs are almost always higher than either BMDLs or BPDLs (Tables II and III; Fig. 3), and therefore may provide less conservative estimates than the other PoD values. They may also be less preferred because of their dependence on the specific doses tested.

As part of this effort to expand the number of data analysis tools for determination of PoDs, other approaches beyond the L&L and BMD approaches were examined and evaluated. The L&L and segmented models provided similar results (Tables II and III). However, the segmented PoD may be viewed as a more reliable metric since it does not require dataset censoring at the highest doses to address saturation or high-dose toxicity, and moreover, it is a well-documented R procedure. The seg-

were combined for simplicity. NOGEL, no observed genotoxic effect level; LOGEL, lowest observed genotoxic effect level; BPD, breakpoint dose; BPDL, breakpoint dose lower confidence interval; STD, slope transition dose; STDL, slope transition dose lower confidence interval; BMD<sub>1SD</sub>, benchmark dose 1 standard deviation, BMDL<sub>1SD</sub>, benchmark dose 1 standard deviation lower confidence interval; BMD<sub>10</sub>, benchmark dose 10; BMDL<sub>10</sub>, benchmark dose 10 lower confidence interval. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]

mented package also contains functions that directly account for variance heterogeneity and non-normality of residuals *via* weights, and conducts these analyses within the framework of generalized linear models (GLMs). GLMs can directly model data sets such as cell or colony counts, incidence frequencies, and other types of data that do not generally adhere to the normality and variance constancy assumptions. Therefore, we propose that the segmented package supersedes the L&L package as the preferred bilinear modeling approach for assessing genetic toxicology data. The STD method, which uses a smoothing regression spline approach also within a GLM framework, directly assesses the slope of the dose-response relationship for a continuous non-linear dose-response function rather than for two linear segments (bilinear model); it provided PoD estimates that were consistent with those derived by other methods (Fig. 3). A potential advantage of the smoothing regression spline approach compared to bilinear modeling, is that it can be compared directly to different non-linear models such as the exponential, Hill and quadratic models. Thus we recommend the smoothing regression spline method applied here in preference to the segmented and L&L models. However, this method shares a major disadvantage similar to the BPD approaches, in that it is frequently not possible to derive a PoD with this method. For example, a PoD was determined for only 15/34 using smoothing regression spline and segmented methods (Table IV), whereas PoDs were determined for 30/34, 31/34, and 34/34 datasets when using NOGEL, BMD<sub>10</sub>, and BMD<sub>1SD</sub>, respectively (Tables II and III).

Similar to observations in the previous evaluations of EMS and MMS data (Gollapudi et al., 2013), the BMD

TABLE IV. Descriptions, Advantages, Disadvantages, and Potential Limitations of the PoD Metrics Examined in This Study

	Full name	Definition	Defined using	Other output metrics	Potential limitations	Advantages
NOGEL	No observed genotoxic effect level	The highest tested dose for which there is no statistically significant increase in genotoxic effect compared to the control.	Dunnett's, Dunnett's Td, Dunn's	LOGEL, <i>P</i> value	Lower power tends to provide larger PoDs; Statistical assumptions must be met; Highly dependent on the study design, e.g., dose selection and dose spacing	Easy to apply; Does not require dose response modeling; Commonly defined. In Tables II and III, it was calculated for 30/34 datasets
BPD	Breakpoint dose	The dose at which the slope changes from zero (horizontal) to positive, with its standard error forming the confidence bounds (90% CI)	L&L	BPDL, BPDU, <i>P</i> value, <i>y</i> intercept, gradient after BPD	Based on one model; Inflexible in terms of ability to account for other functional forms; Ability to define a BPD is highly dependent on the study design, e.g., dose selection and dose spacing. Not commonly defined. In Tables II and III it was calculated for only 8/34 datasets using L&L and 15/34 using segmented, furthermore when the prerequisites were considered, this was reduced to 2/34 and 3/34 respectively.	Lower power tends to provide smaller PoDs; May be appropriate when mechanistic data are available
STD	Slope Transition Dose	The lowest dose for which the lower bound of the 95% confidence interval of the slope exceeds zero	mgcv	STDL, STDU, <i>P</i> value	Plotting of the non-linear model is possibly too flexible compared to pre-defined models; Model is still being developed and validated; Ability to define an STD is highly dependent on the study design, e.g., dose selection and dose spacing. In Tables II and III it was calculated for only 15/34 datasets	Lower power tends to provide smaller PoDs; May be appropriate when mechanistic data are available; Unlike the BPD, can readily be compared to other models e.g. quadratic; Less affected by distribution and variance
BMD <sub>10</sub>	Benchmark Dose 10	A dose that produces a 10% increase over the fitted background	PROAST	BMDL <sub>10</sub> , BMDDU <sub>10</sub> , BMDDU/BMDL ratio	Requires consensus on appropriate biologically relevant benchmark response (BMR); Continuous and quantal data are modeled differently; Often produces very low BMDL metrics	Lower power tends to provide smaller PoDs; Fits function to entire dose-response, not just the tested doses; Currently used by many regulatory agencies; Suitable for use in comparing PoD metrics between different endpoints; Requires fewer data points than BPD and STD; Commonly defined. In Tables II and III it was calculated for 31/34 datasets
BMD <sub>1SD</sub>	Benchmark Dose 1 Standard Deviation	~10% excess risk for individuals below and above the 2nd and 98th percentiles, respectively	BMDS	BMDL <sub>1SD</sub> , BMDDU <sub>1SD</sub> , BMDDU/BMDL ratio	Requires consensus on appropriate biologically relevant benchmark response (BMR); Continuous and quantal data are modeled differently; Comparisons between endpoints and historical datasets more influenced by background level and variance than the BMD <sub>10</sub> approach	Lower power tends to provide smaller PoDs; Fits function to entire dose-response, not just the tested doses; Currently used by many regulatory agencies; Requires fewer data points than BPD and STD; Commonly defined. In Tables II and III it was calculated for 34/34 datasets

NOGEL, no observed genotoxic effect level; BPD, breakpoint dose; BPDL/U, breakpoint dose lower/upper confidence interval; STD, slope transition dose; STDL/U, slope transition dose lower/upper confidence interval; BMD<sub>1SD</sub>, benchmark dose 1 standard deviation; BMDL/U<sub>1SD</sub>, benchmark dose 1 standard deviation lower/upper confidence interval; BMD<sub>10</sub>, benchmark dose 10, BMDL/U<sub>10</sub>, benchmark dose 10 lower/upper confidence interval.

<sup>a</sup>When statistical power is reduced, the PoD is also reduced due to the increased variance in the dataset.

was readily determined for almost all of the ENU and MNU datasets. Moreover, the BMD approach provides a number of advantages when compared to the NOGEL, BPD and STD methods. For example, the BMD methodology generally requires fewer doses in comparison to the BPD methods, BMDLs are readily defined [i.e., Tables II and III, and (Gollapudi et al., 2013)], and BMDL<sub>10</sub> values, although generally lower and thus more conservative, are comparable to other PoDs for the datasets analyzed here. It is worth noting that the BMDL<sub>10</sub> value, which represents a lower confidence limit of a 10% increase above the estimated background, is often below the BPDL, which represents a 0% change above background [Tables II and III and also in (Gollapudi et al., 2013)]. This is a consequence of these two PoD metrics being defined using different statistical models and approaches (Crump, 2011; Slob and Setzer, 2013).

An essential feature of a PoD approach that can be broadly applied with ease and success is its ability to determine whether, and at what dose, there is a detectable increase in genotoxic effect above the spontaneous background level in a particular system. It has been stated that a BPD cannot be accurately defined unless the sample size is infinite (Crump, 2011; Slob and Setzer, 2013); therefore, when examining the low dose region, a PoD metric that is based on a specified increase above a selected background (e.g., the BMD) may be the more relevant approach (Slob and Setzer, 2013). Based on the analyses conducted here, the previous work of Gollapudi et al. (2013), and the work conducted at the RIVM (Hernández et al., 2010, 2011a, b, 2012, 2013; Slob and Setzer, 2013) we support a recommendation to use the BMD approach for assessing dose responses for continuous genetic toxicology data unless otherwise justified. This approach has the added advantage of unifying analyses of genetic toxicology data with analyses of other types of toxicology data.

As part of a related exercise, we employed Monte Carlo simulation to empirically assess the effect of dataset censoring (i.e., varying number of doses and dose spacing) on the probabilistic distribution of PoD values for a given dataset. The results obtained to date, which will be published separately, indicate that BPD determinations (e.g., the L&L BPD) are far more sensitive to dataset censoring than are the BMD determinations. For example, analyses of *in vitro* HPRT gene mutation dose-response data for ENU (Doak et al., 2007) indicated that BPD determination is optimized when the dataset contains responses for three or more doses below the PoD and three or more doses above the PoD. No such requirement was noted for the same dataset with respect to the determination of a BMD<sub>10</sub>. Moreover, although not typically applied, censoring of data near the point of inflection can prevent successful determination of a BPD value. *A priori* assessment of the number of doses most suitable for BMD analysis indicates that three or more doses and

a control are a reasonable starting point, although additional doses will typically improve the precision of the estimated PoD. The precision of the BMD can be indicated by the BMDL to BMDU ratio.

The importance of being able to define a usable PoD metric for all datasets is underscored by the subsequent comparisons of BMDL values across the endpoints investigated. For example, the results presented in Table II suggest that significant increases in *Dlb1* mutations in the small intestine occur at lower ENU doses than that required to elicit significant increases in *LacZ* transgene mutations (Table II). For MNU, significant increases in *Hprt* gene mutations in spleen tissue occur at lower doses in comparison with that required to elicit a significant increase in *LacI* transgene mutations (Table III). These lower PoDs for *Dlb1* and *Hprt* mutation may reflect differences in assay sensitivity, and/or gene target differences (i.e., the ability to discriminate responses in treated from control), and/or differences in repair capacity in the different tissues. However, when focusing on genotoxicity potency ranking across the compounds investigated in this and the earlier Gollapudi et al study (i.e., lowest to highest BMDL<sub>10</sub>) (Table V), these differences have less impact than one might expect; the order of potency *in vitro* and *in vivo* for both gene mutation and MN endpoints are consistent despite variations in strains and endpoints. The ranking of substances from most potent to least potent is MNU > ENU > MMS > EMS for each endpoint based on the BMDL<sub>10</sub> value derived using PROAST. There are very limited carcinogenicity data for these compounds; these include the following: a 54-week MNU mouse study, a 104-week MNU rat study with one dose, a 113-week ENU rat study, and a MMS mouse study with one dose. There are no carcinogenicity studies for EMS (Carcinogenic Potency Database); therefore, it is difficult to compare carcinogenicity rankings. Nevertheless, the lowest genotoxicity BMDL<sub>10</sub> values for ENU and MNU, 0.001 and 0.85 mg/kg/day, respectively, are within two orders of magnitude as the reported TD<sub>50</sub> values for MNU (i.e., 0.0927 and 1.23 mg/kg/day for rats and mice, respectively). Moreover, the TD<sub>50</sub> values reported in the Carcinogenic Potency Database (Table V) yield a ranking from most potent to least potent of MNU > ENU > MMS. This corresponds to the aforementioned ranking based on genetic toxicity BMDs, and provides additional support for the use of genetic toxicology PoDs in human health risk assessment.

#### Incorporating Genetic Toxicity PoD Values into Human Health Risk Assessment

The main focus of this effort was evaluation of several methods for determining genetic toxicity PoDs. However, it is also important to highlight how genetic toxicology PoD metrics can be employed in human health risk

**TABLE V. Table of Lowest BMDL Values Defined Using PROAST for MNU and ENU, Along with Previously Defined PoD Metrics from Gollapudi et al. (2013)**

		MNU	ENU	MMS	EMS
Gene mutation	<i>in vitro</i> (μM)	0.006 <sup>a</sup>	0.68 <sup>a</sup>	4.72 <sup>b</sup>	8.70 <sup>a</sup>
	<i>in vivo</i> (mg/kg)	0.0007 <sup>c</sup>	0.09 <sup>d</sup>	1.34 <sup>e</sup>	9.29 <sup>f</sup>
Micronucleus	<i>in vitro</i> (μM)	0.03 <sup>a</sup>	0.17 <sup>a</sup>	1.00 <sup>g</sup>	4.35 <sup>a</sup>
	<i>in vivo</i> (mg/kg)	0.02 <sup>h</sup>	1.36 <sup>d</sup>	1.74 <sup>i</sup>	56.68 <sup>f</sup>
Cancer bioassay	<i>in vivo</i> (mg/kg/day)	0.093 <sup>j</sup>	0.95 <sup>j</sup>	31.8 <sup>j</sup>	Not available

Order of potency is MNU>ENU>MMS>EMS for genetic toxicology BMDL<sub>10</sub>, and is MNU>ENU>MMS for cancer bioassay TD<sub>50</sub>. Most potent to least potent PoDs are shown from left to right.

<sup>a</sup>Doak et al. (2007), Gene Mutation: *HPRT* gene, AHH-1 cells, 24 hr treatment. Micronucleus: AHH-1 cells, 18 hr treatment.

<sup>b</sup>Pottenger et al. (2009), *Tk* gene, L5178Y cells, 4 hr treatment.

<sup>c</sup>BMS (unpublished data), Rat, *Pig-a* gene, RET and RET cells, 28 days gavage.

<sup>d</sup>van Delft et al. (1998), Mouse, *Dlbl* gene, small intestine, 1 day i.p.

<sup>e</sup>Roche (unpublished data); Rat, *Pig-a* gene, RBC cells, 28 days gavage.

<sup>f</sup>Gocke and Wall (2009), Gene Mutation: *LacZ* gene, Mutagenome, bone marrow cells, 28 days gavage. Micronucleus: bone marrow cells, 7 days gavage.

<sup>g</sup>Bryce et al. (2010), TK6 cells, 24–30 hr treatment.

<sup>h</sup>LeBaron (2009), Rat, Blood, 4 days gavage.

<sup>i</sup>LeBaron et al. (2008), Rat, Blood, 4 days gavage.

<sup>j</sup>Lowest TD<sub>50</sub> from the Carcinogenic Potency Database (<http://toxnet.nlm.nih.gov/cpdb/>). Values adjusted for differences in treatment duration.

assessment, e.g., to support determination of regulatory limits to reduce or to quantify the risk of adverse genotoxic effects in humans. We introduce the potential role of such PoDs in risk assessment briefly, while acknowledging that comprehensive recommendations on quantitative approaches for the use of genetic toxicity data in regulatory decision making will require additional analyses and discussion. In this regard, we note that the analyses and recommendations reported herein and by Gollapudi et al. (2013) have been considered by the recent Working Group on Quantitative Approaches to Genetic Toxicology Risk Assessment that met as part of the International Working Groups on Genotoxicity Testing (IWGT) in Brazil in November 2013 (IWGT, 2013). This group is preparing two publications that endorse many of our recommendations and provide additional recommendations for the use of PoD metrics in human health risk assessment.

#### **Mode of Action (MOA) Data to Support Extrapolation Below the PoD**

It is necessary to have chemical-specific MOA information to justify the assumption of different slopes in the dose–response curve below the PoD as compared to above the PoD. Conversely, the dose–response analyses can also be used to support the MOA information. For example, recent work by Johnson and colleagues has shown that DNA repair capabilities below the PoDs serve to counteract specific gene mutation and chromosomal damage induced by the alkylating agents MMS, EMS, MNU and ENU (Zair et al., 2011; Johnson et al., 2012; Thomas et al., 2013). The data reported by Zair et al., (2011) support a role for the DNA repair enzyme methyl-

purine DNA glycosylase (*MPG*) in repair of clastogenic lesions below the PoD; when *MPG* levels were reduced by RNA interference, the EMS PoD shifted to a sevenfold lower concentration (i.e., BMDL<sub>10</sub> decreased from 1.19 μg/mL to 0.17 μg/mL). A similar decrease in PoD was shown with the methyl-guanine methyl-transferase (*MGMT*) DNA repair activity and the mutagenic lesion O<sup>6</sup>-alkylG, where prior *MGMT* inactivation using the nucleotide analogue O<sup>6</sup>-benzyl guanine reduced the MNU PoD for *HPRT* mutant frequency to an approximately 50-fold lower concentration (Thomas et al., 2013). Moreover, over-expression of *MGMT* was shown to significantly protect against, but not completely nullify, the effect of MNU in tumor initiation (Becker et al., 2013). Such data provide support for a biological mechanism underlying a non-linear dose response in the region around the PoD. Those studies focused on EMS and MNU, respectively, for clastogenicity and mutagenicity, but the similarity in the types of DNA adducts and mutation spectra for EMS, MMS, ENU, and MNU (Beranek, 1990; Jenkins et al., 2005; Jenkins et al., 2010; Sharm et al., 2014) suggests that this group of mono-functional alkylating agents have efficient DNA repair mechanisms operating below the PoD that dramatically diminish their genotoxic effects. These cellular processes provide a mechanism for differences in the dose–response slope below and above the PoD.

#### **Using BMDL<sub>10</sub> to Support Regulatory Evaluation**

PoD metrics from toxicology endpoints are frequently used to support the determination of regulatory limits that can be employed to manage the risk of adverse health

effects. We suggest the use of PoDs from *in vivo* genetic toxicology endpoints, in conjunction with, or in some circumstances in place of, PoDs estimated for other observed adverse effects. For example, the BMD<sub>10</sub> PoD values for ENU and MNU defined in this study (Tables II and III), revealed lowest *in vivo* BMD<sub>10</sub> values for ENU and MNU of 0.09 (male mice, *Dlb-I<sup>a/b</sup>* gene mutation) (van Delft et al., 1998) and 0.0007 mg/kg/day (male rat Charles River Crl:CD (SD), RET *Pig-a* gene mutation) (BMS *Pig-a* RET Day 29), respectively. If the BMD<sub>10</sub> doses from the *in vivo* assays are converted to human equivalent doses by using the respective scaling factors of 0.081 for mouse to human, and 0.16 from rat to human (FDA, 2005), 7.3 and 0.11 µg/kg/day, or 438 and 6.6 µg/day for a 60 kg human, are obtained as the human equivalent doses associated with the aforementioned rodent BMD<sub>10</sub> values. These PoDs can be used in a similar manner to PoDs from other toxicity endpoints, e.g., to determine a regulatory limit such as a reference dose (RfD) after application of suitable uncertainty/safety factors. If, for example, a conservative safety factor of 100 (i.e., 10× for animal to human extrapolation and 10× for variability in human populations) is applied to the above PoDs, a calculated tolerable daily intake based on this endpoint would be 4.37 and 0.07 µg/person/day for ENU and MNU, respectively. If circumstances warranted, there could be a reason for the uncertainty factor related to human variability to be reduced (e.g., from 10 to 3) or even removed, based on the ability of the BMD approach to account for variability in the data as compared to, for example, the NOGEL. For example, a study with lower statistical power and greater variance would produce a lower BMD (and BPD/STD), but a higher NOGEL. If chemical-specific adjustment factors (CSAFs) are available for interspecies differences and human variability, their use also would be considered in the selection of uncertainty/safety factors (WHO/IPCS, 2005). Therefore, when considering methods for incorporation of PoD metrics into human health risk assessments, it is important to remember that some PoD metrics are more conservative than others. Moreover, the same uncertainty/safety factors would not necessarily be applicable when using the BMD compared to the NOGEL, and thus should not necessarily be applied in all instances.

A related approach that also uses the genetic toxicology PoDs estimated above involves the calculation of the increasingly used margin of exposure (MOE) metric. This approach is becoming preferable, as it incorporates estimated or actual human exposure information in the overall assessment. It is a straightforward method that involves comparison of the PoD and the current or predicted human exposure (i.e., a simple ratio of the PoD to human exposure). Regulatory decision making, and the requirement for risk management interventions, are based on the magnitude of the ratio; a larger MOE is less of a

concern (e.g., MOE ≥ 10,000 may be considered to present minimal risk), while a smaller MOE may be less acceptable (e.g., MOE < 100). Other considerations that help determine the “acceptability” of the MOE approach include the severity of the effect, the MOA, the number of adverse effects observed, whether the observed effect(s) are from animal or human studies, the number of assumptions used in MOE estimations, the size of the affected population, and whether any susceptible subgroups have been identified.

Whether one uses the RfD approach or an MOE approach for genetic toxicity endpoints, the results can be evaluated with all the other available toxicity data to provide an improved and informed human health risk assessment. If determination of the PoD, and the subsequent comparative assessment indicates that genetic toxicity is the driving concern for human health considerations, then genetic toxicity data could become the basis for regulatory decision making.

## CONCLUSIONS

1. MNU and ENU both elicit sub-linear dose responses that yield PoD metrics for gene mutation and chromosomal damage endpoints *in vitro* and *in vivo*.
2. Among the methods/approaches investigated here, the BMD approach yields the most conservative PoDs (i.e., BMD<sub>10</sub>).
3. The BMD<sub>10</sub> is comparable to, and recommended alongside, the BMD<sub>ISD</sub>, as the most suitable metrics for defining PoDs for continuous genetic toxicology data.
4. The BMD method is the preferred PoD determination method, followed by the NOGEL method, the smoothing regression spline to determine the STD, and then the segmented methods to determine the BPD (now supersedes the L&L for BPD determination).
5. PoD metrics from genetic toxicology dose-response data, *via* the derivation of regulatory limits such as the RfD or risk management metrics such as the MOE, can be used for evaluations of human risk and regulatory decision making.

Routine determination of PoD metrics for genetic toxicity dose-response data, and routine use of genotoxicity PoD values for regulatory evaluations of new and existing substances will require application of the preferred methodology (i.e., the BMD approach) to a wider range of compounds with a diverse array of MOAs. We have already begun collecting and analyzing detailed dose-response data for other recognized genetic toxicants. Comparative analyses of the PoD values across a variety of endpoints, including carcinogenicity, as well as estimates of regulatory limits analogous to RfDs where appropriate, will enhance the foundation for the routine

interpretation of genetic toxicity dose-response data in a human health context.

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## AUTHOR CONTRIBUTIONS

Drs. George Johnson, Bhaskar Gollapudi and Paul White provided leadership for the GTTC QAW during database construction, data collection, and data analysis, and led discussions of the group on results interpretation as well as preparation of the manuscript. Dr. Johnson is also the lead author of the paper. Drs. Greg Hixon, Chad Thompson and Liz Abraham developed and implemented algorithms for analyses of the dose-response data and determination of point of departure metrics. Drs. George Johnson and Lya Soeteman-Hernández also participated in the analyses of the dose-response data. Drs. Lya Soeteman-Hernández, Owen Bodger, Kerry Dearfield, Robert Heflich, Greg Hixon, David Lovell, Jim MacGregor, Lynn Pottenger, Chad Thompson, Liz Abraham, Veronique Thybaud, Jan van Benthem and Errol Zeiger provided insightful review of the data analyses, the data analysis methodologies and the interpretation of the results, and participated in preparation of the manuscript. Dr. Jennifer Young Tanir provided logistical, organizational and editorial support for the project. All authors contributed to addressing the comments from the internal reviewers.

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